



Experimental and modeling study of visible light responsive photocatalytic oxidation (PCO) materials for toluene degradation

Lexuan Zhong^a, James J. Brancho^b, Stuart Batterman^{a,*}, Bart M. Bartlett^b, Christopher Godwin^a

^a Environmental Health Sciences, University of Michigan, Ann Arbor, MI, USA

^b Chemistry, University of Michigan, Ann Arbor, MI, USA

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ABSTRACT

Only limited research has examined the development and application of visible light responsive photocatalytic oxidation (PCO), although such materials have great potential for mitigating concentrations of volatile organic compounds (VOCs) when applied to building surfaces. This study evaluates the performance and characteristics of a visible light responsive photocatalyst, specifically, a co-alloyed TiNbON compound with a band energy of 2.3 eV. The PCO material was developed using urea-glass synthesis, characterized by scanning electron microscopy (SEM), diffuse reflectance spectra (DRS), powder X-ray diffraction (PXRD), and Brunauer-Emmett-Teller (BET) methods, and VOC removal efficiency was measured under visible light for toluene (1–5 ppm) at room temperature (21.5 °C) over a range of relative humidity (RH: 25–65%), flow rate (0.78–7.84 cm/s), and irradiance (42–95 W/m²). A systematic parametric evaluation of kinetic parameters was conducted. In addition, we compared TiNbON with a commercial TiO₂-based material under black light, estimated TiNbON's long-term durability and stability, and tested its ability to thermally regenerate. Using mass transfer and kinetic analysis, three different Langmuir-Hinshelwood (LH) type reaction rate expressions were proposed and evaluated. A LH model considering one active site and competitive sorption of toluene and water was superior to others. The visible-light driven catalyst was able to remove up to 58% of the toluene, generated less formaldehyde than the commercial TiO₂, could be fully regenerated at 150 °C, and had reasonable durability and stability. This evaluation of TiNbON shows the potential to remove VOCs and improve air quality for indoor applications. Further research is needed to evaluate the potential for harmful by-products, to identify optimal conditions, and to use field tests to show real-world performance.

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1. Introduction

Indoor and outdoor air pollution represents a challenging public health issue that is receiving increasing attention in both developed and developing countries. Poor air quality in China and India, for example, reduces average life expectancy by an estimated two years, and globally, air pollution is considered to be the fourth-largest environmental threat to human health [1]. Volatile organic compounds (VOCs), which are precursors to ozone (O₃) and particulate matter (PM), play a critical role in air quality. VOCs are emitted from motor vehicles, chemical manufacturing facilities, refineries, factories, consumer and commercial products, and biogenic sources, among numerous other sources [2–7]. Pollutants

in outdoor air also affect indoor air quality (IAQ). While filtration and air cleaning in a building's heating, ventilating and air conditioning (HVAC) system can improve IAQ, filters installed in typical HVAC systems control only PM and do not remove VOCs [8]. Gas phase filtration of VOCs is expensive and applications have been limited [9]. Moreover, many buildings do not utilize mechanical HVAC systems and thus have limited opportunities for filter-based air cleaning systems. These challenges have motivated the development of innovative technologies that can control gaseous pollutants and improve IAQ.

Photocatalytic oxidation (PCO) technology is an emerging advanced technology that promises offers new and cost-effective opportunities for improving IAQ. PCO materials can be applied on indoor surfaces, including walls, windows, floors and ceilings, and potentially can utilize visible light provided by room lighting and sunlight to oxidize VOCs. These materials could represent a sustainable or "green" approach if effective visible-light driven catalysts can be developed. While some PCO-based technologies have been

* Corresponding author at: M6075 SPH II, 1415 Washington Heights, Ann Arbor, MI 48109-2029, USA.

E-mail address: stuartb@umich.edu (S. Batterman).

evaluated [10–19], research addressing the development and evaluation of visible light driven PCO materials is limited [20–23]. It is known, however, that materials such as TiO₂ can be co-alloyed with niobium (Nb) and oxynitrites to significantly reduce adsorption energy down to 2.0 eV, which enables visible light activation [24–26]. Most evaluations have focused on NO_x (NO, NO₂) removal using UV lights for outdoor applications [27–32]. Several studies have evaluated VOC removal (e.g., ethanol, heptane, decane, benzene, toluene, *p*-xylene, and acetone) at relatively high concentrations, e.g., tens to hundreds of ppm [33–38], but these levels are well above those usually encountered outside relatively few industrial environments. Other studies have evaluated VOC removal using heated PCO reactors [39,40], which are not suitable for indoor applications and which have other cost and environmental issues, and to date, field studies of commercial PCO-based products have not demonstrated VOC removal [41,42]. Hence, the development of reliable and efficient PCO materials for remains a research priority.

An important concern with PCO as well as other VOC removal and treatment technologies is the formation of hazardous by-products [43]. For example, degradation of benzene by O₃-assisted PCO has been shown to form octane, decane, nonane, decanal, and decanoic acid (all in gas-phase), as well as (solid phase) *p*-benzoquinone [43]. PCO degradation of toluene formed benzene, and degradation of *p*-xylene produced both benzene and toluene [38]. Under simulated solar irradiation, TiO₂-based paints exposed to *n*-decane formed C6-C11 alkanes, 4-methylnonane, 2,6-dimethyloctane, butanoic acid, propanoic acid and butanal [44]. Formaldehyde was the major by-product in several PCO studies [18,45–48]; this compound is of special concern as it is a potent human carcinogen. Another concern is that PCO materials tested at low VOC concentrations (e.g., ppm levels) can produce byproduct levels too low for quantification by many gas chromatography (GC) methods [49]. Because PCO mechanisms are complex and experimental conditions in the literature vary considerably, it is difficult to compare by-product formation.

The stability and durability of PCO coatings on building substrates is another key issue. PCO materials must be long-lived for practical applications, and thus materials must both resist abrasion and maintain chemical activity. While considerable effort has been devoted to improving photocatalytic activity, few studies have examined PCO stability on surfaces [12]. Accelerated weathering (mostly rain rinsing) and aging tests have been conducted for TiO₂-coated materials [50–53], however, such tests are designed for outdoor applications. Tests of abrasion resistance for TiO₂-based building materials in indoor applications have remained scarcely explored.

The objective of this paper is to evaluate the performance of a new PCO material that could be applied to building surfaces to improve air quality. We characterize this material, present the design of a system intended to evaluate PCO performance, show results pertaining to toluene removal using visible light-driven TiO₂-based films, and evaluate the formation of formaldehyde and other byproducts using sensitive GC-MS methods. In addition, we examine long-term durability and physical stability of the coatings using a simple but novel peeling-off test, applied for the first time (to our knowledge) to evaluate resistance to mechanical abrasion. Lastly, we fit the experimental data using mass transfer and kinetic models, which also help to explain and extend results.

2. Experimental

2.1. Materials and chemicals

A new TiNbON catalyst was prepared by co-alloying TiO₂ with Nb and oxynitrites using urea-glass synthesis in which TiCl₄ reacted

with NbCl₅ and urea in an ethanol solution. The urea-glass reaction was modified using an additional oxidation step to prepare the active catalyst (details pending publication elsewhere). As a reference PCO material, we used Aerioxide P25 TiO₂ (80% anatase, 20% rutile, BET area = 47.39 m²/g, Evonik Degussa GmbH, Essen, Germany). Aluminum plates (85 mm × 105 mm × 0.2 mm) were used as substrates; plates were cleaned to remove potential surface impurities using acetone followed by methanol rinses. The test gas used to evaluate PCO removals was prepared in filtered air (see below) using reagent grade toluene (99.9%, Sigma-Aldrich, St. Louis, USA).

2.2. Film preparation using the doctor-blade technique

To prepare the reference PCO material, 500 mg of P25 TiO₂ was added to a plastic cup, along with zirconia mixing balls and 2.7 mL *n*-methyl pyrrolidone (NMP). NMP was selected as a solvent (rather than water or polymer additions) based on preliminary tests of PCO performance and its (see Supplemental Figs. S1 and S2). The cup was placed in a planetary mixer (Thinky Mixer AR-100, Laguna Hills, USA) and mixed for 10 min to create a homogeneous and stable suspension. Then, 0.8 mL of the suspension was spread onto one of the substrates using the doctor blade technique, then the substrate was placed in a drying oven at 160 °C for 30 min to evaporate the NMP. A total of 20 substrate plates was similarly prepared and used for these tests. These substrates, denoted as P25N, had a catalyst loading of 0.5 ± 0.03 mg/cm², determined as the mass difference before and after coating divided by the area coated. The same approach was used to prepare 40 substrates, each using 500 mg of TiNbON; these plates had a loading of 1.2 ± 0.5 mg/cm².

2.3. Catalyst characterization

The catalyst surface structure and particle sizes were characterized by scanning electron microscopy (SEM; Physical Electronics Auger Nanoprobe 680, Physical Electronics, Inc., Eden Prairie, USA) using a 10 kV beam voltage and 10 nA beam current. Nitrogen sorption isotherms were measured at 77.3 K using a Quantachrome NOVA 4200 analyzer (Quantachrome Instruments, Boynton Beach, USA) and the Brunauer-Emmett-Teller (BET) method to calculate the specific surface area. Diffuse reflectance spectra (DRS) were collected on a UV-vis-NIR spectrophotometer (Cary 5000, Agilent Technologies, Santa Clara, USA) using a Harrick Praying Mantis diffuse reflectance accessory and BaSO₄ as a 100% reflectance standard. Reflectance data were modified for presentation in Tauc plots (Fig. 3, inset) by the indirect Kubelka-Munk function. The crystallite structure of TiNbON was examined using powder X-ray diffraction and Cu K α radiation (PXRD, D8 Advance, Bruker, Daltonics Inc., Billerica, USA).

2.4. PCO test reactor setup

A schematic diagram of the experimental set-up is shown in Fig. 1. The total air flow (5–8 L/min) of VOC-free air at 25, 50 or 65% relative humidity (RH), which spans the RH comfort range [54], was prepared by blending carbon-filtered dry and water saturated (using bubblers) flows at appropriate ratios as controlled by calibrated rotameters (DryCal flow meter, Brandt Instruments, USA). VOC concentrations in the range of 1–5 ppm were generated by injecting neat toluene into the air flow at a constant rate of 1.4–10.5 μ L/h (depending on the concentration and air flow) using a 25 μ L syringe and a syringe pump with accuracy of ±0.5% and reproducibility of ±0.1% (Model 74900, Cole Parmer Instrument, Vernon Hills, USA). Tubing, fittings and most other materials in contact with the toluene flow were Teflon and stainless steel.

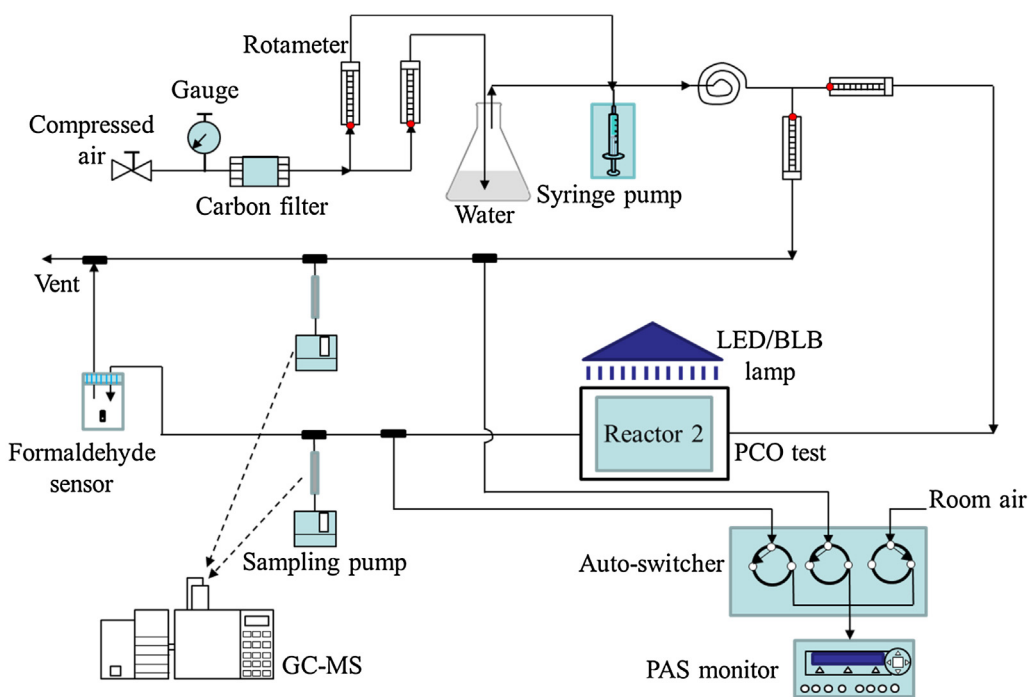


Fig. 1. Schematic diagram of the PCO system. Only 1 reactor is shown.

PCO substrates were tested using two identical aluminum reactors with internal volumes of 0.12 L. The reactors were gas-tight and equipped with a removable glass window and fittings for incoming and existing flows. Substrates were placed in the center of the reactor at a depth of 5.0 ± 0.5 mm below the glass window. PCO efficiencies were measured using test gas flow rates from 0.2 to 2 L/min. Flows measured before and after each test varied by less than 1%. Given the reactor dimensions and flow rates, test flows of 0.2, 1.0 and 2.0 were in the laminar regime ($Re = 5, 25, 50$ from Eq. (7)), and the surface velocity across the substrate was 0.78, 3.92 and 7.84 cm/s, respectively. A 12 W LED flood lamp with a temperature color of 5000 K (Model 9290011835, Philips, Somerset, USA) placed 5.0–12.5 cm from the PCO substrate gave an irradiance of 42–95 W/m² (400 – 540 nm) as measured by a radiometer (Solarimeter, Solartech Inc., Harrison Charter Township, USA). We also tested a 75 W fluorescent black light (BL) using an irradiance of 7.5 ± 0.4 W/m² (365 nm).

VOC concentrations during each test were measured nearly continuously using a photoacoustic spectrometer (PAS, INNOVA 1312, California Analytical Instruments, Orange, USA) and periodically using a gas chromatograph/mass spectrometer (GC/MS, Model 6890/5973, Agilent Technologies, Santa Clara, USA). The PAS was coupled to a 3-channel automatic switcher that alternately selected reactor inlet and outlet flows at 1 min intervals, thus providing a near-continuous record of concentration changes. Due to the PAS's high air sampling flow rate (1.3–2 L/min) and lack of VOC speciation, VOC levels for the PCO reactions were quantified by GC/MS. Duplicate inlet and outlet samples were collected on adsorbent tubes containing 160 mg 60/80 mesh Tenax-GR by pumping for 2–5 min at 2.5 ± 0.1 mL/min (low flow module, Aircheck Sampler, SKC Inc., Pittsburgh, USA); flows were determined using a bubble meter. Prior to use, the adsorbent tubes were conditioned in a flow of ultrapure nitrogen at 325 °C for 6 h. Adsorbent samples were injected onto the capillary column of the GC/MS using an automated short-path thermal desorption/cryofocusing system (Model 2000, Scientific Instrument Services, Ringoes, USA), which measures a wide range of VOCs with excellent quantification for VOCs eluting between pentane and naphthalene [55]. GC/MS parameters

are reported in Supplemental Table S1. Formaldehyde at the reactor exit was measured by photoelectric photometry (GrayWolf FM-801, GrayWolf Sensing Solutions, Shelton, USA) using 0.5 h samples in both PCO and regeneration modes.

The PAS was calibrated using toluene at concentrations of 1, 2, 3, 5 and 10 ppm. Method detection limits (MDL) were determined as the standard deviation of seven replicate low concentration injections multiplied by 3.14. MDLs were 0.25 ppm (942 µg/m³) for the PAS, 0.02 ppb (0.07 µg/m³) for toluene using the GC/MS, and 4.88 ppb (6 µg/m³) for formaldehyde.

2.5. Test protocol

Initial tests were conducted to optimize the system, confirm that it was leak-tight, that sink effects were negligible (<0.5% change between inlet and exit concentrations), and that the air supply was clean (no detectable VOCs). Flow stability was confirmed. Additional tests were conducted using PCO-coated substrates to determine appropriate VOC sampling volumes and flow rates, and to investigate the presence of by-products.

Prior to each test, a new substrate was placed in the reactor, and the reactor system was cleaned by purging for 1 h with clean air. Then, the desired humidity and VOC levels were set and maintained for 1–2 h to condition and stabilize reactor conditions, as confirmed by real-time VOC (PAS) and humidity measurements. Supplemental Table S2 details the many experimental conditions used to examine UV-PCO performance. Prior to lamp illumination, dark adsorption tests were conducted. Once the PAS confirmed steady-state conditions, the PCO reaction was initiated by turning on the visible lamp, maintained for 1 h (6 h in stabilization tests, see below) so that outlet concentrations were stable (supported by Supplemental Fig. S3), duplicate VOC samples collected at reactor inlet and outlet ports to characterize PCO efficiency and by-products, and formaldehyde was measured for 0.5 h in the outlet flow. After these tests were completed, the toluene injection was stopped and the lamp kept on to regenerate the photocatalytic activity on the catalyst surface. Formaldehyde also was measured during regeneration.

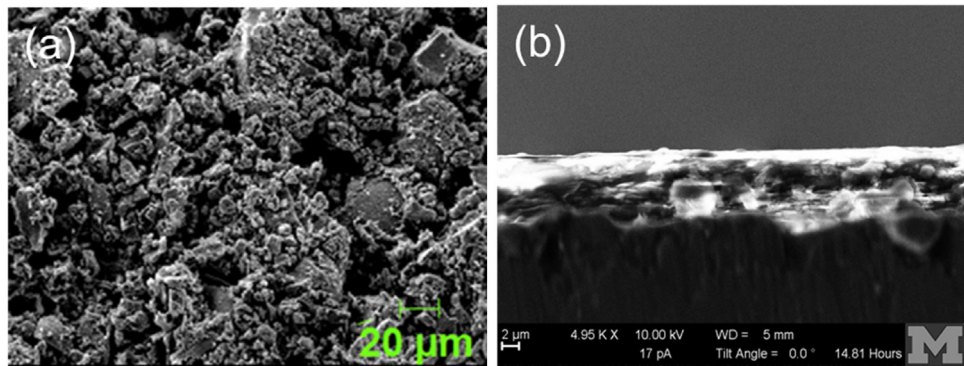


Fig. 2. SEM micrographs of TiNbON: (a) top view, and (b) side view.

2.6. Stability

The stability of the coated substrate was evaluated in two aspects: mechanical stability establishing the integrity and adhesion of the catalyst on the substrate; and the catalyst or photocatalytic stability regarding VOC removal efficiency. A “peel-off” test evaluated mechanical stability. This entailed cutting a 19 × 25 mm section of tape (3 M United States, St. Paul, USA), weighing it on a microbalance (1 µg resolution), placing the sticky side of the tape on the catalyst surface, gently pressing it against the substrate, and then peeling off the tape at 2.5 or 0.25 cm/s to characterize adhesion at two abrasion levels. The tape was weighed a second time to determine the amount of PCO that became detached. This test was repeated five times at each peeling speed (using new tape strips) on randomly selected but unused sections of the substrate.

PCO stability was determined using toluene and five exposure cycles with the same substrate, each cycle consisting of 6 h of exposure to 1 ppm toluene at 95 W/m². Duplicate VOC samples were taken every 2 h. In order to understand any potential new chemical bonds formed after PCO exposure, Raman spectra for the coating before and after PCO exposure was collected on a inVia Raman spectrometer (Renishaw Inc., West Dundee, USA) equipped with a Leica microscope, RenCam CCD Detector, a 785 nm diode laser, and 1200 lines/mm grating, and Raman data was analyzed using the Renishaw's WiRE software package. An earlier study demonstrated that ultraviolet C (UVC)-induced regeneration can partially restore the photocatalytic activity, however, recovery was limited after long-term UVC exposure [56]. We investigated the ability to increase PCO lifetime and speedup recovery and regeneration of PCO-based materials using a thermal method in which the substrate was heated to 150 °C for 0.5 h in an oven. A second set of five exposure cycles was conducted using the same conditions except that the substrate was thermally treated between each cycle.

2.7. Quantification metrics

The single-pass removal efficiency E (%) represents the percentage of pollutant removed from the air stream after passing through the PCO reactor:

$$E(\%) = \frac{Q(C_{in} - C_{out})}{QC_{in}} \times 100 = \frac{C_{in} - C_{out}}{C_{in}} \times 100 \quad (1)$$

where C_{in} (mg/m³) = upstream contaminant concentration at steady state; C_{out} (mg/m³) = downstream contaminant concentration at steady state, and Q (L/min) = flow rate through the reactor. This definition of efficiency is used throughout this paper.

The surface reaction and by-product generation rates are based on the amount of pollutant removed and amount of by-product

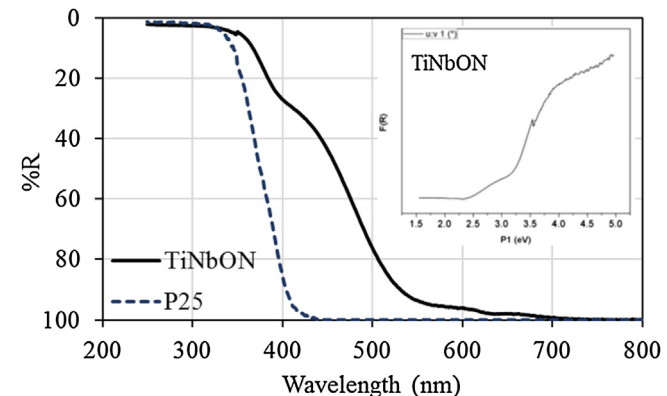


Fig. 3. Diffuse reflectance spectra of TiNbON and P25 catalysts.

generated, respectively, on the catalysts effective BET surface area per unit time at steady state:

$$-r_{PCO,ave} = \frac{Q(C_{out} - C_{in})}{aA_{BET}m} \quad (2)$$

$$r_{bp,ave} = \frac{Q(C_{out} - C_{in})}{aA_{BET}m} \quad (3)$$

where A_{BET} = BET surface area of catalyst (m²/g), m = catalyst mass on the tested piece (g), and a = constant accounting for effective catalyst exposure to light, here a value of 0.8 was considered for evaluation on the assumption of exponential decay of the irradiance distribution within the catalyst layer thickness [17]. Another way to estimate the effective illuminated surface area uses the average aggregate size and average effective light path [57]. In this study, the assumed value of $a = 0.8$ is justified by the comparison between experimental measurements and simulation results, which is presented in Section 3.5.

3. Results and discussion

3.1. Powder/film characterization

TiNbON is a fine bright yellow powder. SEM micrographs in top and side views (Fig. 2) show nanoparticles that agglomerated into irregularly shaped surfaces. Aggregates were several hundred nm to several µm in size, and the primary particles had a mean diameter of approximate 1–2 µm. The BET surface area was 40.76 m²/g, which indicates that TiNbON was as porous as P25 (50 m²/g), despite large differences in particle morphology. XRD indicated that the TiNbON phase composition was 100% in the anatase phase (Supplemental Fig. S4), different from P25 which was 80% anatase and 20% rutile. The film thickness averaged 4–5 µm (Fig. 2b).

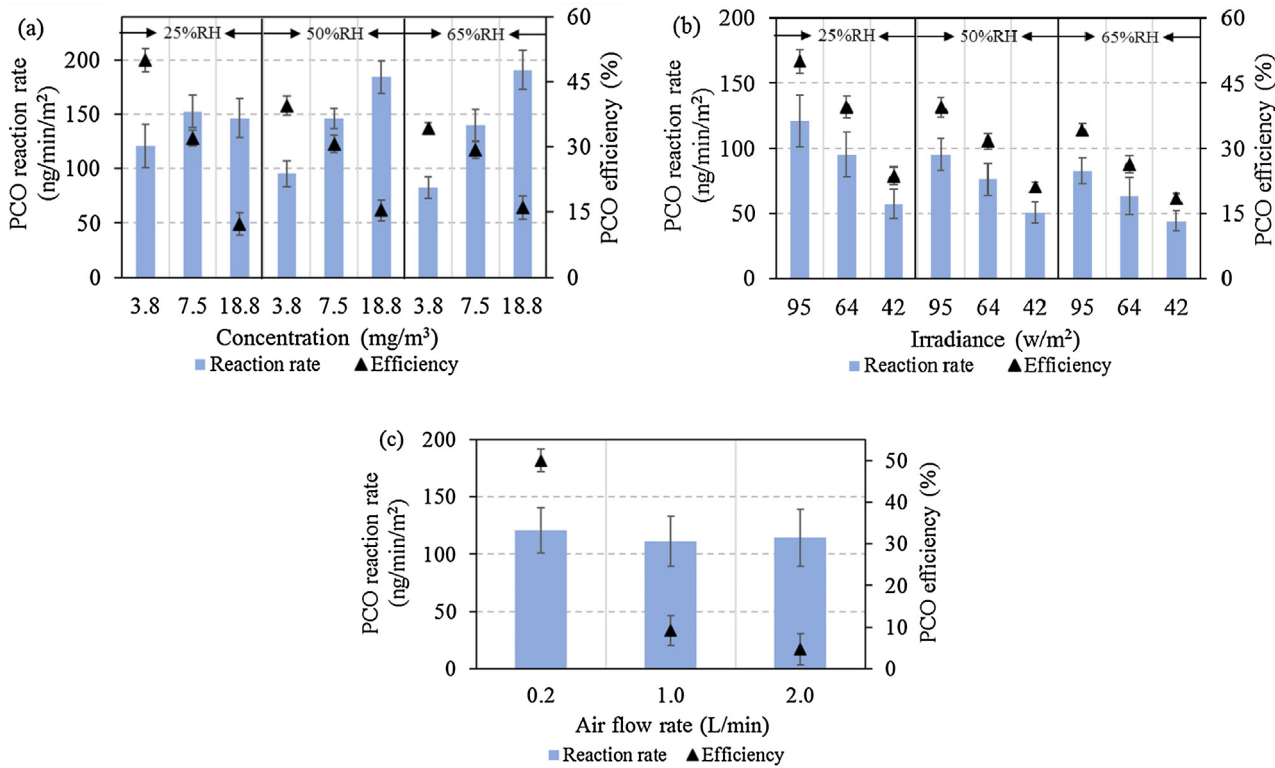


Fig. 4. Influence of operation conditions on the PCO of toluene: (a) Effect of concentration and humidity ($I=95\text{ W/m}^2$); (b) Effect of irradiance and humidity ($C_{\text{in}}=3.8\text{ mg/m}^3$); and (c) Effect of air flow ($C_{\text{in}}=3.8\text{ mg/m}^3, I=95\text{ W/m}^2$).

The UV-vis diffuse reflectance spectra of TiNbON and P25 (Fig. 3) show that most adsorption spectra of P25 were located below 400 nm, and that the absorption edge of TiNbON red-shifts from UV into the visible region (500–600 nm). This provides evidence that Nb and N co-alloyed into the TiO_2 lattice. The bandgap of TiNbON was estimated as 2.3 eV, corresponding to visible light at 540 nm.

3.2. Effects of operating parameters on toluene removals

Operating parameters affected PCO performance in the steady-state tests using toluene. Fig. 4a shows the effect of humidity and toluene inlet concentration. The single-pass PCO removal efficiency consistently decreased as toluene concentrations increased from 3.8 to 18.8 mg/m³; this applied at each humidity (25, 50 and 65% RH). This can be attributed to the interaction of toluene and water molecules adsorbed on the catalyst surface: given the same humidity, higher concentrations increase the number of toluene molecules per unit time entering the reactor, and a higher number of hydroxyl radicals (OH^-) would be required to achieve the same efficiency, however, the amount of catalysts on the substrate surface is fixed, which potentially limits OH^- production given the constant RH and irradiance conditions. Hence, efficiencies decrease at higher concentrations, as shown previously [33]. The PCO reaction rate data reveal some seemingly inconsistent trends, e.g., the highest rate occurred at 7.5 mg/m³ and 25% RH, and the apparent rate increased with inlet concentration at both 50% and 65% RH. Fig. 4a also indicates that reaction rates depended strongly on humidity. For inlet concentrations of 3.8 and 7.5 mg/m³, the efficiency and PCO reaction rate decreased as RH increased from 25 to 65%. However, for an inlet concentration of 18.8 mg/m³, humidity slightly increased PCO efficiency and reaction rates. These results can be explained by existence of an “optimal” humidity level at each toluene concentration: water is needed for the PCO process since it provides the required OH^- for oxidation (i.e., a shortage

of water molecules will lower performance due to lack of OH^-), however, at high humidity, competitive adsorption may exclude toluene from the substrate and thus inhibit PCO conversion. For example, moderate humidity, then dry conditions were shown to provide the highest reaction rates in the oxidation of benzene at high concentrations (100 ppm) [37]. An “optimal” humidity level can balance the inhibition of physical adsorption with the promotion of photochemical conversion [18]. These effects are explored in Section 3.4 and 3.5.

PCO efficiency and reaction rate were proportional to the level of irradiance (Fig. 4b), which generates electron-hole pairs that initiate PCO reactions by reacting with water to form OH^- radicals. Again, RH levels affected results. More electron-hole pairs on the catalyst surface are produced at higher irradiance, however, recombination of photo-generated charge carriers can reduce the overall quantum efficiency [58]. The recombination time scale has been estimated to range from 1 to 10 picoseconds, much shorter than the time scale of charge carriers trapped by surface chemicals (nanoseconds) [59]. Hence, enhanced irradiance can be effectively utilized for the PCO reaction.

The effect of flow rate is shown in Fig. 4c. While PCO efficiency decreased by 10-fold as air flow rates increased from 0.2 to 2 L/min, PCO reaction rates were nearly constant. This indicates that the surface photochemical reaction rate, and not the mass transfer between gas and solid phases in the reactor, was the controlling step, i.e., the higher VOC loadings at the higher flow rate decreased removal efficiency given the similar reaction rate on the surface. Although the residence time in the reactor was reduced from 13 to 1.3 s as air flow rates increased from 0.2 to 2.0 L/min, residence times remained much longer than the PCO reaction time (ns). Again, this indicates that surface PCO reactions were the controlling process.

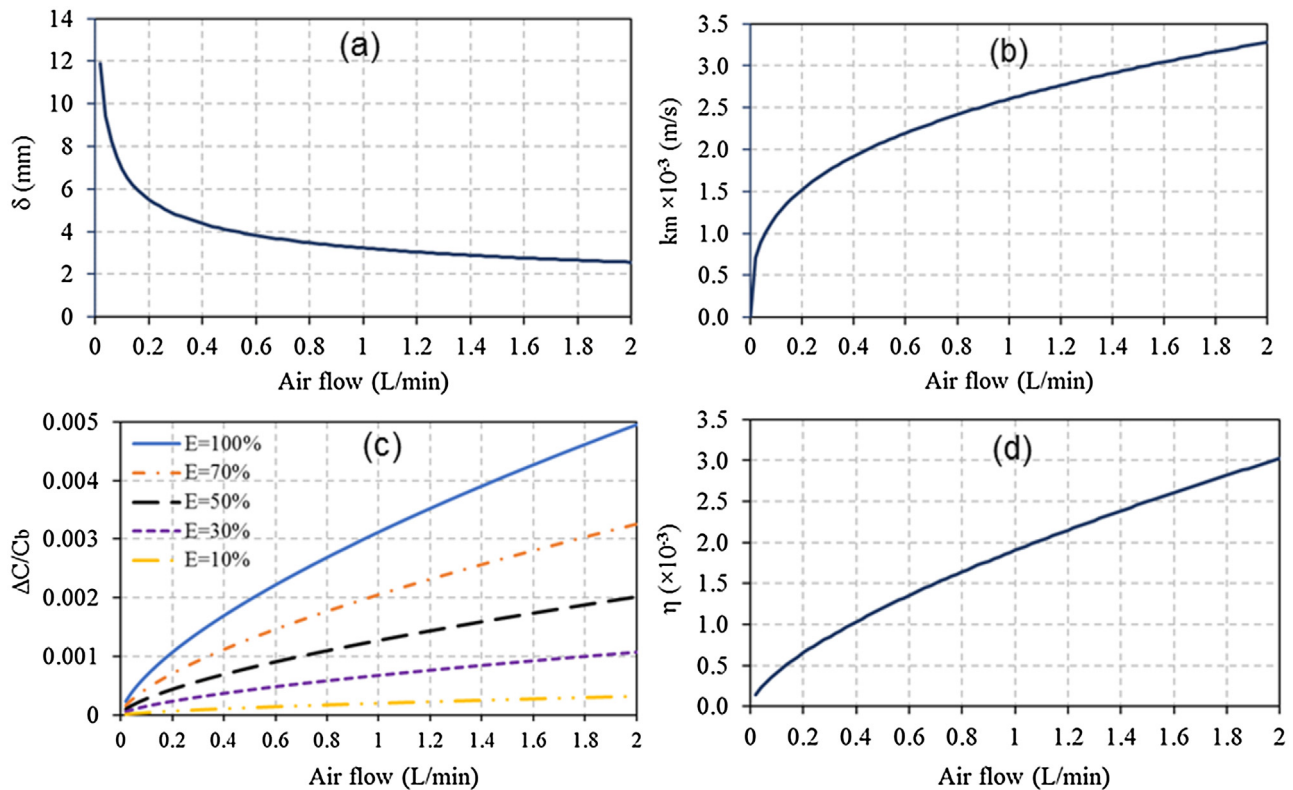


Fig. 5. Influence of air flow in convective mass transfer over PCO materials on: (a) concentration boundary layer thickness; (b) mass transfer coefficient; (c) concentration gradient; and (d) η .

3.3. By-product formation

During the PCO of toluene, no detectable other gas-phase by-products other than formaldehyde were detected in the reactor outlet flow. One possible reason is our use of low concentrations: most studies aimed at observing by-products have used test gases with much higher VOC levels, e.g., hundreds of ppm [33–38]. A second possible reason is the small volume sampled on the adsorbent tube (selected so as not to overload the GC/MS with toluene) that may have limited sensitivity for the generated by-products (possible in the pg range). In addition, some of the anticipated by-products (e.g., methanol, acetaldehyde, benzaldehyde, acetic acid [14,60]) are unlikely to be trapped effectively and detected by the adsorbent sampling and GC/MS method used. While additional and potentially toxic by-products may be formed by the PCO of toluene, the presence of by-products at very low concentrations may pose insignificant health risks; however, further analyses using different and possibly more sensitive methods are recommended.

3.4. External mass transfer in the reactor

We first assume plug flow (reactor velocity u in m/s), which leads to a concentration boundary layer along the surface (depicted in Supplemental Fig. S5). The mass balance at steady-state for the two phases is:

$$\text{Gas phase : } \varepsilon D \frac{d^2 C_b}{dx^2} - u \frac{dC_b}{dx} - k_m S (C_b - C_s) = 0 \quad (4)$$

$$\text{Solid phase : } k_m (C_b - C_s) = -r_{\text{PCO}} \quad (5)$$

where S = effective BET surface area of catalyst on the plate per unit reactor volume (m^{-1}), C_b = gaseous VOC concentration in the bulk (gas-phase) (mg/m^3), C_s = VOC concentration on the catalyst surfaces (mg/m^3), and k_m = average mass transfer coefficient (m/s). For

laminar flow over a channel, k_m can be estimated from the following empirical correlation for the Sherwood number over the length of a plate, L (0.105 m in our case) [61]:

$$Sh = \frac{k_m d_h}{D} = 1.85 \left(Re Sc \frac{d_h}{L} \right)^{1/3} \quad (6)$$

$$Re = d_h u / \nu \quad (7)$$

$$Sc = \frac{\nu}{D} \quad (8)$$

where ν = kinematic viscosity of air for a dilute gas ($15.1 \times 10^{-6} \text{ m}^2/\text{s}$ at 21°C), d_h = hydraulic diameter (m), and D = molecular diffusion coefficient of the VOC, here $8.4 \times 10^{-6} \text{ m}^2/\text{s}$ at 21°C for toluene using the WL method [17]. The boundary layer thickness δ and mass transfer coefficient k_m can be expressed as a function of velocity:

$$\delta = \frac{D}{k_m} = \frac{L^{1/3} d_h^{1/3} D^{1/3}}{1.85 u^{1/3}} \quad (9)$$

$$k_m = \frac{1.85 u^{1/3} D^{2/3}}{(d_h L)^{1/3}} \quad (10)$$

Fig. 5a demonstrates how boundary layer thickness decreases with air velocity, which agrees with previous findings [62]. This is an intuitive result: increasing velocity leads to steeper concentration gradients. The boundary layer thickness always exceeded the height of the shallow (5 mm) PCO reactor. Hence, concentration gradients were large in the vertical direction. Given the slow surface velocity (0.78–7.84 cm/s), the gas required a long time (1.3–13.4 s) to pass over the plate, providing sufficient time for mass transfer from bulk flow to the catalyst surface. **Fig. 5b** indicates that the higher air flows tested increased the mass transfer coefficient.

Table 1
Fitted parameters in the kinetic model.

Model	Competition type	Expression	n	k ng/(min m ²)/(w/m ²) ⁿ	K ₁ (m ³ /mg)	K ₂ (m ³ /mg)	K ₃ (m ³ /mg)	K ₄ (m ³ /mg)	R ²
LH-1	Two types of non-competitive sites	$r = I^n \frac{kK_1C_S}{1+K_1C_S} \frac{K_2C_W}{1+K_2C_W}$	0.8	4.512	1.240	1.612×10^{-3}	–	–	0.814
LH-2	One type of competitive sites	$r = I^n \frac{kK_1C_SC_W}{(1+K_1C_S+K_2C_W)^2}$	0.8	20.625	0.650	1.061×10^{-3}	–	–	0.953
LH-3	Two types of competitive sites	$r = I^n \frac{kK_1C_S}{1+K_1C_S+K_2C_W} \frac{K_4C_W}{1+K_3C_S+K_4C_W}$	0.8	0.0110	–0.389	2.469×10^{-5}	–0.034	-8.587×10^{-5}	–

Assuming that the bulk concentration C_b changes linearly over the plate length in a laminar flow, a reasonable estimate is:

$$C_b \approx \frac{C_{in} + C_{out}}{2} \quad (11)$$

Combining Eqs. (2) and (5),

$$\Delta C = (C_b - C_s) = \frac{Q(C_{in} - C_{out})}{k_m a A_{BET} m} \quad (12)$$

$$\frac{\Delta C}{C_b} = \frac{2Q(C_{in} - C_{out})}{k_m a A_{BET} m(C_{in} + C_{out})} \quad (13)$$

where h=channel height (0.005 m), and w=width of the plate (0.085 m). Eq. (13) further examines the significance of the external mass transfer on the heterogeneous PCO. Normally, ratios greater than 0.15 indicate that the effect of mass transfer is significant [39]. Fig. 5c shows the correlation between air flow and $\Delta C/C_b$ for different PCO efficiency profiles. At a fixed air flow and higher concentration gradient, higher removal efficiency can be reached. If the concentration gradient is fixed, a higher air flow will decrease the PCO removal efficiency. For increases in both the concentration gradient and the air flow, the PCO performance will likely remain the same. For all cases, $\Delta C/C_b < 0.15$, which indicates the external mass transfer is not significant in the reactor for the tested air flow rates (0.2–2 L/min).

$$C_s = C_b - \frac{Q(C_{in} - C_{out})}{k_m a A_{BET} m} \approx \frac{C_{in} + C_{out}}{2} - \eta(C_{in} - C_{out}) \quad (14)$$

$$\eta = \frac{Q}{k_m a A_{BET} m} \quad (15)$$

Fig. 5d shows η as a function of low air flow rates. Eq. (15) can be simplified to:

$$C_s \approx C_b \approx \frac{C_{in} + C_{out}}{2} \quad (16)$$

Thus, the surface concentration is approximately equal to the gaseous concentration, which further supports that external mass transfer has only a negligible influence, again supporting findings in Fig. 5c. Hence, the rate-limiting step is the PCO reaction on the catalyst surface.

3.5. Kinetic study and simulation

Three different Langmuir-Hinshelwood (LH)-based reaction rate expressions accounting for various active sites and competi-

tion were used to describe the surface reaction. The experimental kinetic data were fitted using MATLAB R2016a (MathWork Inc., Natick, USA) and information in Table 1. Several adsorption equilibrium parameters in the LH-3 model yielded negative values, which had no meaning and were thus excluded. Results in Supplemental Fig. 6 show that the LH-2 model ($R^2 = 0.953$) that considers one type of competitive sites was superior to the LH-1 model ($R^2 = 0.689$) that considers two types of non-competitive sites. In addition, the LH-2 predictions almost fell on the 1:1 line with observations (slope = 0.964). This means that the model with one type of bimolecular competitive site provided the most suitable description of this study's experimental data. In contrast, a different "best" model was reported for PCO of n-hexane in a tube reactor [33], possibly due to the different reactor configuration, photocatalysts, challenge gases, or other parameters.

The LH-2 model was employed for simulating the PCO of toluene to examine the influence of a wide range of operational parameters, e.g., RH, irradiance, and inlet concentration, on the PCO reaction rates. C_s in the LH-2 kinetic reaction rate expression (Table 1) is an important parameter, yet it is not forward determined since the outlet concentration is unknown. This problem was solved by determining the correlation between the outlet concentration and one variable (RH or I or C_{in}) when other variables were kept constant. Table 2 lists outlet concentration expressions for three scenarios. Fig. 6a shows that the PCO reaction rate increases rapidly with humidity level, reaches a maximum rate at the "optimal" humidity, and then slowly declines at higher humidities. This applies at each inlet concentration. In addition, the optimal humidity increases with the inlet concentration. In consequence, relatively dry levels will be favorable for the PCO of VOCs in most non-industrial buildings due to the normally low concentrations (ppb level) of VOCs present. Interestingly, PCO reaction rates of toluene at 7.5 mg/m³ are higher than those at 18.8 mg/m³ for RH below 30%, which explains findings in Fig. 4a at 25% RH. These simulations demonstrate the importance of that competitive sorption between water and toluene molecules on the catalyst surface. Similarly, when the humidity and irradiance are constant, an optimal inlet concentration will maximize the PCO reaction rate (Fig. 6c). The impact of irradiation on the PCO reaction rate is nearly linear (Fig. 6b). In general, our experimental measurements agree with these simulations, indicating the LH-2 model describes PCO behavior in this study.

Table 2
Surface concentration expressions.

Conditions	C _{out} (mg/m ³)	R ²	Conditions	C _{out} (mg/m ³)	R ²	Conditions	C _{out} (mg/m ³)	R ²	
I ^a = 95 C _{in} ^b = 3.8 C _{in} ^b = 7.5 C _{in} ^b = 18.8	C _{out} = 6.93 × 10 ⁻⁵ RH + 1.61 C _{out} = 2.64 × 10 ⁻⁵ RH + 4.97 C _{out} = −9.78 × 10 ⁻⁵ RH + 16.9	0.963 0.980 0.939	C _{in} ^b = 3.8 RH = 25% RH = 50% RH = 65%	C _{out} = 20.05I ^{-0.518} C _{out} = 10.12I ^{-0.326} C _{out} = 10.08I ^{-0.313}	0.999 0.999 0.982	RH = 25% RH = 25% RH = 25%	I ^a = 95 I ^a = 64 I ^a = 42	C _{out} = 0.982C _{in} − 2.016 C _{out} = 0.990C _{in} − 1.531 C _{out} = 0.983C _{in} − 0.892	0.999 1.000 1.000

^a Unit of I is W/m².

^b Unit of C_{in} is mg/m³.

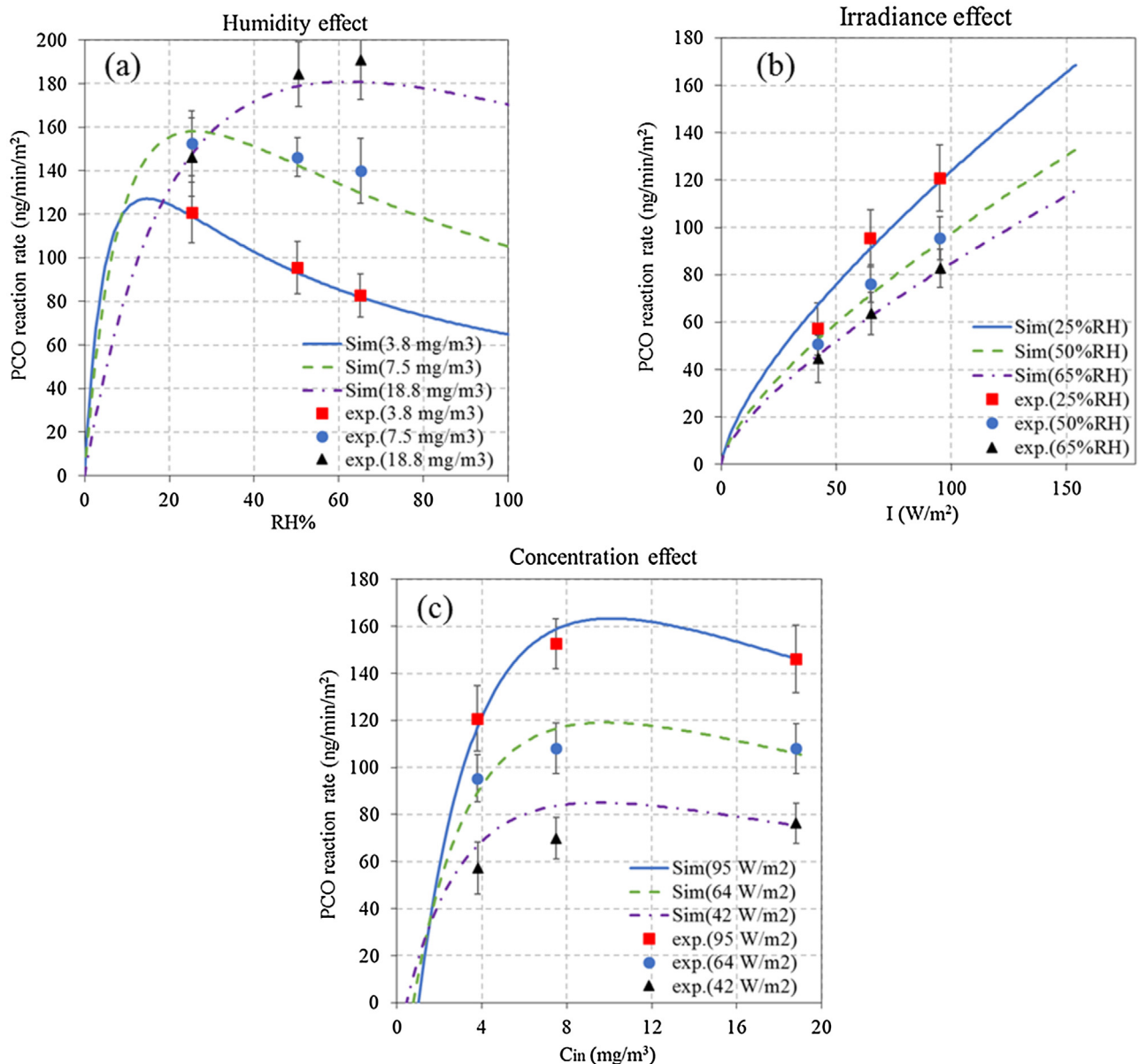


Fig. 6. Simulated and observed effects on the PCO reaction rates for toluene at steady-state: (a) influence of humidity with $I = 95 \text{ W/m}^2$; (b) influence of irradiance with $C_{\text{in}} = 3.8 \text{ mg/m}^3$; and (c) influence of inlet concentration with $\text{RH} = 25\%$.

3.6. Comparison with P25N

PCO performance of TiNbON initiated by a visible light was compared with that of P25N under black light to demonstrate new features of TiNbON that may relate to real world applications. Fig. 7 shows that the PCO reaction rates and formaldehyde generation rates of the two catalysts under several conditions. Usually, formaldehyde generation rates from P25N were 3–5 times higher than those from TiNbON, while PCO reaction rates of P25N were 1–2 fold higher than TiNbON. P25's higher PCO activity may be attributed to the rutile phase (20%) that promotes electron-hole pair separation, thus suppressing recombination [63]. However, P25 had lower ability to mineralize toluene and the by-products increased health risk. It is interesting to observe that P25N was susceptible to RH (from $261 \text{ ng/min}\cdot\text{m}^2$ at 25% RH to $135 \text{ ng/min}\cdot\text{m}^2$ at 65% RH), but TiNbON had slightly lower PCO reaction rate with humidity. Possibly, Nb and oxynitrides may modify the hydrophilicity, rendering TiNbON less vulnerable to humidity, suggesting that TiNbON might apply to a wide range of environments.

In general, operation parameters produced similar trends for both P25N and TiNbON since they were TiO_2 -based catalysts. PCO performance of P25 under visible LED light was negligible (PCO efficiency <2%), implying that TiNbON successfully expands the adsorption spectrum from UV to visible range and has ability to be applied in the building material surfaces.

3.7. Adhesion

The peeling-off test showed average losses of $88 \pm 12 \mu\text{g}/\text{cm}^2$ (loss percentage of $7.9 \pm 1.1\%$) and $222 \pm 20 \mu\text{g}/\text{cm}^2$ ($19.9 \pm 1.8\%$) at peeling-off speeds of 0.25 cm/s and 2.5 cm/s , respectively. These results suggest potentially reasonable resistance to mechanical abrasion. This is further supported a lack of visible particles lost from the coated substrates and left in the reactor throughout the tests.

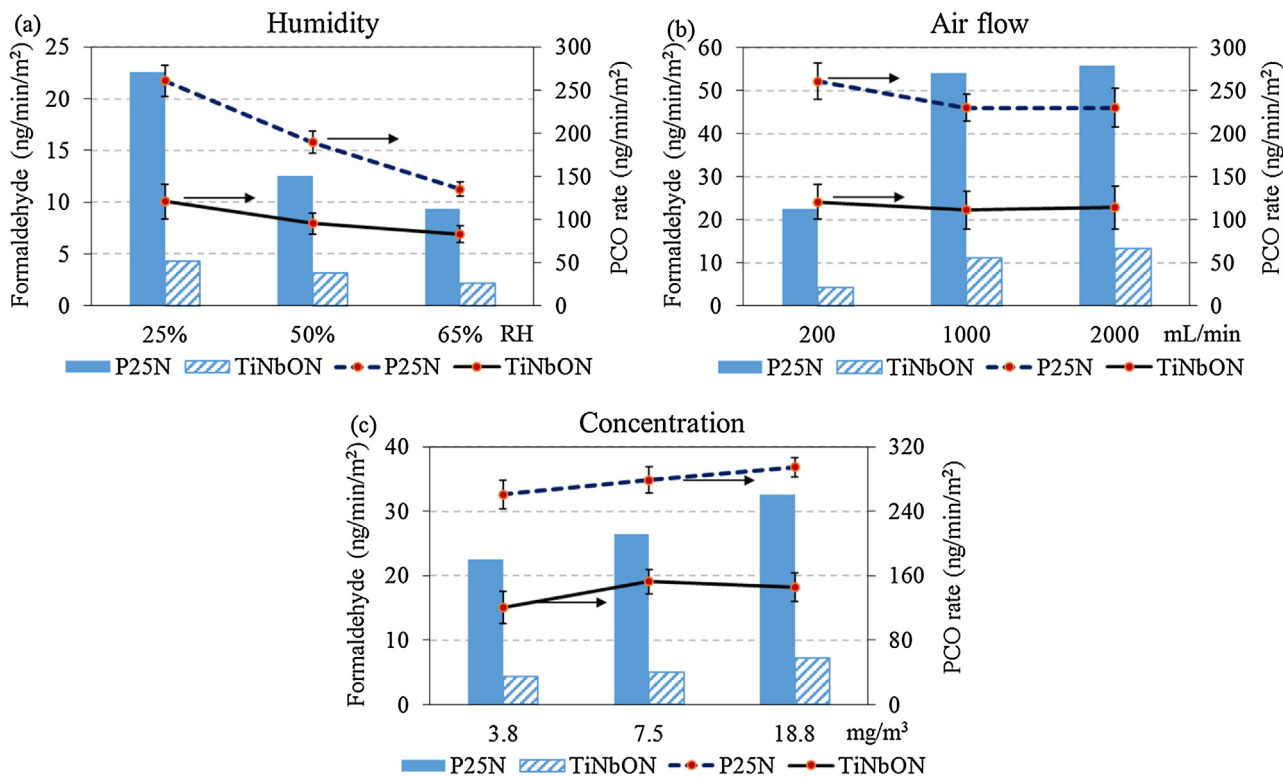


Fig. 7. Comparison of PCO performance for P25N with BLB (7.5 W/m²) and TiNbON with LED (95 W/m²) under varying (a) humidity levels (toluene 3.8 mg/m³, air flow 0.2 L/min), (b) air flow rates (toluene 3.8 mg/m³, 25%RH), and (c) inlet concentrations (25%RH, air flow 0.2 L/min).

3.8. Stability and regeneration

As the amount of chemisorbed-intermediates, e.g., carboxylic acids [14,60], on active sites increased, the net transfer from toluene to intermediates declined and decreased the available vacant sites. In turn, this decreased site availability caused a gradual decline in the adsorption of toluene from the gas-phase, which led to an obvious decline of the PCO reaction rate. After two cycles of photo-degradation, TiNbON exhibited a significant loss of PCO activity, and as the reaction proceeded, reaction rates continued to decrease (Fig. 8) as did generation of the by-product formaldehyde. This can be attributed to chemisorbed intermediates with high affinity that gradually blocked active PCO sites and inhibited water molecules from adsorbing and converting intermediates into small gaseous molecules. We also observed a color change from bright yellow to dark yellow after five cycles. Unfortunately, the Raman results show little difference between the sample after 5-cycle exposure to toluene and the reference. No signals in the organic region (500–2000 cm⁻¹) of the vibrational spectrum were detected. If there was anything by-product adsorbed, it was below the detection limits ($\sim 0.1\text{--}1\text{ mg/cm}^2$) of the measurement. Considering the tested concentration (1 ppm) was higher than the (ppb) levels normally encountered in non-industrial buildings, the deactivation time in real world settings would be much longer than shown in our tests.

The regeneration performance shown in Fig. 8 demonstrates that thermal conditioning maintained PCO rates at $119 \pm 2\text{ ng/min/m}^2$ without significant change over the five repeated cycles. Thus, thermal regeneration effectively removed the refractory intermediates and fully restored the PCO activity.

4. Conclusion

This study characterized the performance of a co-alloyed TiNbON compound, a visible light active photocatalytic (PCO) material that can oxidize toluene and other VOCs. Compared to other TiO₂-based materials, the band gap of TiNbON was reduced to 2.3 eV with the responsive edge wavelength of 540 nm, and the removal efficiency of toluene under visible light was high. A parametric evaluation of TiNbON performance used toluene concentrations from 1 to 5 ppm, relative humidities from 25 to 65%, air velocities from 0.78 to 7.84 cm/s, and irradiances from 42 to 95 W/m². PCO efficiency increased at lower air velocities and higher irradiance levels, and interactions with humidity and inlet concentrations could be explained using a Langmuir-Hinshelwood-based kinetic model using one active site with competition of toluene and water molecules at the catalyst surface. The surface PCO reaction, rather than mass transfer, was the rate-limiting step in the reactor. Formaldehyde was detected as a by-product of both the PCO and regeneration processes. In comparison to other TiO₂-based materials, TiNbON had greater ability to resist the influence of humidity and generated lower levels of formaldehyde. Finally, the long-term durability tests show that TiNbON gradually lost PCO activity, but thermal regeneration could restore its performance, and that the material had reasonable abrasion resistance.

The experimental results suggest that TiNbON films applied to building materials may represent a promising option to reduce VOC pollutants in indoor applications. PCO materials can work indoors, and the ongoing transition of lighting technologies in buildings (from incandescent, to fluorescent and now to LEDs) allow successful application of PCO technology. However, further research is needed to evaluate the potential for harmful by-products, to identify optimal conditions and means to regenerate PCO materials, and to use field tests to show real-world performance.

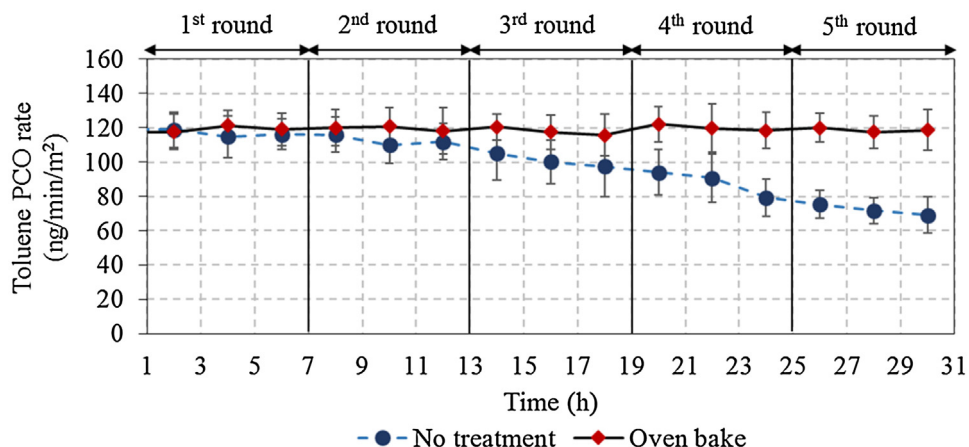


Fig. 8. Stability of TiNbON for toluene removal after 5 6-h test cycles (toluene 3.8 mg/m³, 25%RH, 0.2 L/min, LED 95 W/m²).

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Appendix A. Supplementary data

Supplementary data associated with this article can be found, in the online version, at <http://dx.doi.org/10.1016/j.apcatb.2017.05.047>.

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